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(54) Titre : PROCEDE DE PREPARATION D'UNE FORMULATION DE PROTEINE THERAPEUTIQUE ET
FORMULATION D'ANTICORPS PRODUITE PAR UN TEL PROCEDE

(54) Title: METHOD OF PREPARING A THERAPEUTIC PROTEIN FORMULATION AND ANTIBODY FORMULATION
PRODUCED BY SUCH A METHOD

(57) Abrégé/Abstract:

The invention relates to a method of preparing a protein formulation including a therapeutic protein, the method comprising the steps of: providing a solution comprising said protein; concentrating the protein in the solution by a first ultra-filtration step; diafiltrating the solution with a diafiltration buffer including at least one first excipient, whereby a retentate is obtained comprising the protein and the first excipient; further concentrating the protein in the retentate by a second ultra-filtration step; and adding at least one final excipient, whereby the protein formulation with a desired protein concentration is obtained. According to the invention, the method further comprises, before the second ultra-filtration step, adding a second excipient to the retentate obtained from the diafiltration step. The invention is also directed to antibody formulations produced by the foregoing method.

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(54) Title: METHOD OF PREPARING A THERAPEUTIC PROTEIN FORMULATION AND ANTIBODY FORMULATION PRODUCED BY SUCH A METHOD

(57) Abstract: The invention relates to a method of preparing a protein formulation including a therapeutic protein, the method comprising the steps of: providing a solution comprising said protein; concentrating the protein in the solution by a first ultra-filtration step; diafiltering the solution with a diafiltration buffer including at least one first excipient, whereby a retentate is obtained comprising the protein and the first excipient; further concentrating the protein in the retentate by a second ultra-filtration step; and adding at least one final excipient, whereby the protein formulation with a desired protein concentration is obtained. According to the invention, the method further comprises, before the second ultra-filtration step, adding a second excipient to the retentate obtained from the diafiltration step. The invention is also directed to antibody formulations produced by the foregoing method.

**METHOD OF PREPARING A THERAPEUTIC PROTEIN FORMULATION AND
ANTIBODY FORMULATION
PRODUCED BY SUCH A METHOD**

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Technical field

10 The present invention relates to a method of preparation of a protein formulation including excipients and at least one therapeutic protein.

The invention is of particular interest in the field of antibody formulations intended for a therapeutic use and is also directed to an antibody formulation produced by the method.

15

Background of the invention

The invention is more particularly related to methods sequentially comprising:

- providing a solution comprising said protein;
- concentrating the protein in the solution by a first ultra-filtration step;
- diafiltering the solution thus obtained with a diafiltration buffer including at least one first excipient, whereby a retentate is obtained comprising the protein and the first excipient;
- further concentrating the protein in the retentate by a second ultra-filtration step in an ultra-filtration equipment;
- adding at least one final excipient, whereby the protein formulation with a desired protein concentration and including said first and final excipients is obtained.

30 In general, the final protein formulations for therapeutic antibodies include at least an amino-acid, such as histidine, which is added during the diafiltration step, and a sugar acting as a stabilizer, such as trehalose. The trehalose is commonly added with the other excipients in the final addition step.

In conventional methods applied to therapeutic antibodies, the above steps are performed with a protein solution, once purified by a number of purification steps usually including a virus retaining filtration as the last purification step. The protein

5 solution (or "product") is concentrated by the first ultra-filtration step, from a concentration of approximately 5 to 20 g/l to a concentration of about 40 to 100 g/l (depending on the protein). Then the concentrated product is diafiltered in a diafiltration buffer, such as histidine. In some instances, the diafiltration buffer may be another standard buffer such as acetate, tris or phosphate. The diafiltration buffer

10 is chosen based on the final protein formulation as well as on any offset that is required due to the Donnan effect. The Donnan effect occurs as the product is concentrated and results in the exclusion of certain charged buffer species, e.g. histidine. The diafiltration buffer is therefore usually adjusted to a higher buffer concentration and a lower pH than are specified for the protein formulation. Once

15 the diafiltration is complete, the product goes through the second ultra-filtration step for concentration to approximately 50% above the desired protein concentration for the final protein formulation. Then the product is removed from the ultra-filtration system and the system is rinsed to recover additional product. With a final concentration of more than 50% above the desired concentration in the protein

20 formulation, all of the rinse can be added back to the product to maximize recovery, without excessively diluting the product. Then the excipients are added (sugar, surfactant, chelator, etc.) as a concentrated solution, usually with a dilution ratio of approximately 4, meaning that 1 unitary volume of the concentrated excipient solution is added to 3 unitary volumes of the product. The dilution ratio of 4 is based

25 on the maximum solubility of the sugar component of the excipient solution, which is usually the limiting factor. If necessary, the product is then further diluted with formulation buffer for adjustment to the final desired concentration.

Such conventional methods may therefore not be applicable when it is desired to

30 obtain a highly concentrated protein in the final formulation, and even more when the protein is of particularly high viscosity.

For example, in the case of a therapeutic antibody formulation with a desired final concentration of 150 g/l, the viscosity of the molecule precludes concentrating to the targeted value of 50% above the desired final concentration.

5

It is an aim of the invention to provide a method of preparation of a protein formulation that may be applied to highly viscous and highly concentrated proteins.

It is a further aim of the invention that the method may be implemented at a 10 manufacturing scale, without negatively affecting the overall yield of the manufacturing process and without incurring extra costs due to an excessive waste of certain excipients. In particular, it is an aim to keep the use of the sugar components, which are particularly costly, at a similar level as the conventional methods.

15

It is still a further aim to preserve the stability of the protein over all the steps of the method and to protect the protein from aggregation.

20 **Summary**

According to a first aspect of the present invention, there is provided a method of the above type further comprising, before the second ultra-filtration step, adding a second excipient to the retentate obtained from the diafiltration step.

25

By moving the addition of a second excipient, in particular the trehalose (or more generally the sugar), to post-diafiltration, the remaining excipients can be added at a much higher concentration in a subsequent step, thus generating a lower dilution of the product. This in turn means that the maximal required concentration can be 30 brought to only about 10% (in some instances between 5 to 15 %) above the final desired concentration, as compared to the value of about 50% for the conventional methods. This 10% value is obtainable with standard ultra-filtration equipments, even with higher molecule viscosity. This also allows recovering product from a rinse and thus allows obtaining a 90% yield of the ultra-filtration/dia-filtration process.

Also, adding the second excipient (the sugar) before the final concentration protects the protein from aggregation.

5 According to preferred embodiments of the invention:

- the method further includes, after step (e) and before step (f), rinsing the ultra-filtration equipment with a rinse buffer, whereby the recovery of the protein is enhanced;
- 10 - the rinse buffer comprises the first and the second excipients at concentrations substantially equal to, respectively, the concentrations of the first and of the second excipients in the protein formulation;
- the first excipient is an amino-acid, preferably histidine;
- the first excipient in the protein formulation has a concentration of between 16 and 24 mM, preferably of between 17 and 23 mM, most preferably of about 20 mM;
- 15 - the second excipient is a sugar, preferably a disaccharide;
- the final excipients include a surfactant, preferably polysorbate 80;
- the final excipients include a chelating agent, preferably EDTA;
- 20 - the protein formulation has a protein concentration of between 110 and 165 g/l;
- the protein is an antibody.

In a first preferred embodiment:

25

- the antibody is an anti-PCSK9 (Proprotein Convertase Subtilisin Kexin type 9) antibody;
- the anti-PCSK9 antibody is selected from the group consisting of bococizumab, evolocumab (REPATHATM), alirocumab (PRALUENTTM), REGN728, 31H4, 11F1, 12H11, 8A1, 8A3, 3C4, 300N, 1D05, LGT209, RG7652, and LY3015014;
- 30 - the anti-PCSK9 antibody comprises a heavy chain variable region (VH) comprising complementarity determining region one CDR1, CDR2, and CDR3 of the amino acid sequence shown in SEQ ID NO: 1; and a light chain

variable region (VL) comprising CDR1, CDR2, and CDR3 of the amino acid sequence shown in SEQ ID NO: 2; or alternatively the anti-PCSK9 antibody comprises a VH CDR1 having the amino acid sequence shown in SEQ ID NO: 3, 4, or 5, a VH CDR2 having the amino acid sequence shown in SEQ ID NO: 6 or 7, a VH CDR3 having the amino acid sequence shown in SEQ ID NO: 8, a VL CDR1 having the amino acid sequence shown in SEQ ID NO: 9, a VL CDR2 having the amino acid sequence shown in SEQ ID NO: 10, and a VL CDR3 having the amino acid sequence shown in SEQ ID NO: 11;

5

10 - the protein formulation has a protein concentration of between 135 and 165 g/l, preferably of between 142 and 158 g/l, most preferably of about 150 g/l;

- the second excipient in the protein formulation is trehalose at a concentration of between 67.2 and 100.8 g/l, preferably of between 71.4 and 96.6 g/l, most preferably of about 84 g/l;

15 - the final excipients include polysorbate 80 which, in the protein formulation, has a concentration of between 0.16 and 0.24 g/l, preferably of between 0.17 and 0.23 g/l, most preferably of about 0.2 g/l;

- the final excipients include EDTA which, in the protein formulation, has a concentration of between 0.04 and 0.06 g/l, preferably of between 0.0425 and 0.0575 g/l, most preferably of about 0.05 g/l;

20 - the protein formulation has a pH of between 5.2 and 5.8, preferably of about 5.5;

- the solution provided in step (a) has a protein concentration of between 5 and 20 g/l;

25 - the protein is concentrated to between 80 and 120 g/l, preferably to between 90 and 110 g/l, and most preferably to about 100 g/l, by the first ultra-filtration step;

- the protein is concentrated to between 143 and 173 g/l, preferably to between 150 and 166 g/l, and most preferably to about 158 g/l, by the second ultra-filtration step;

30 - the first excipient in the diafiltration buffer has a concentration higher than the concentration of the first excipient in the protein formulation, said concentration of the first excipient in the diafiltration buffer being preferably of between 29.75 and 40.25 mM, most preferably of about 35 mM;

- the diafiltration buffer has a pH of between 5.1 and 5.5, preferably about 5.3;
- adding the second excipient to the retentate obtained from the diafiltration step is achieved by adding a first additive solution to the retentate, said first additive solution comprising the second excipient at a concentration of between 340 and 460 g/l, preferably of between 380 and 420 g/l, most preferably of about 400 g/l;
- the first additive solution comprises the first excipient at a concentration lower than the concentration of the first excipient in the diafiltration buffer and higher than the concentration of the first excipient in the protein formulation, said concentration of the first excipient in the first additive solution being preferably of between 25.5 and 34.5 mM, most preferably of about 30 mM;
- the first additive solution further comprises a final excipient;
- the first additive solution comprises about 30 mM histidine and about 400 g/l trehalose;
- adding the first additive solution to the retentate is performed at a dilution ratio of about 4.15, whereby one volume of the first additive solution is added to approximately 3.15 fold the same volume of the retentate;
- adding the final excipients includes the step of adding a second additive solution to the solution obtained from the second ultra-filtration step, said second additive solution comprising the second excipient at a concentration lower than the concentration of the second excipient in the first additive solution and higher than the concentration of the second excipient in the protein formulation;
- the second additive solution comprises the first excipient at a concentration substantially equal to the concentration of the first excipient in the protein formulation;
- the second additive solution comprises about 20 mM histidine, about 84 g/l trehalose, about 1 g/l EDTA and about 4 g/l polysorbate 80;
- adding the second additive solution is performed at a dilution ratio of about 20, whereby one volume of the second additive solution is added to approximately 19 fold the same volume of to the solution obtained from the second ultra-filtration step.

In a second preferred embodiment:

- the antibody is an anti-IL7R antibody;
- 5 - preferably, the anti-IL-7R antibody comprises a heavy chain variable region (VH) comprising complementarity determining region one CDR1, CDR2, and CDR3 of the amino acid sequence shown in SEQ ID NO: 13 (examples of the sequences of such CDRs are SEQ ID NOs. 17, 18 and 19 respectively); and a light chain variable region (VL) comprising CDR1, CDR2, and CDR3 of the amino acid sequence shown in SEQ ID NO: 14 (examples of the sequences of such CDRs are SEQ ID Nos. 20, 21 and 22 respectively);
- 10 - more preferably, the VH region of the anti-IL-7R antibody comprises the amino acid sequence shown in SEQ ID NO. 13, and the VL region of the anti-IL-7R antibody comprises the amino acid sequence shown in SEQ ID NO. 14;
- 15 - even more preferably, the heavy chain of the anti IL-7R antibody comprises the amino acid sequence shown in SEQ ID NO. 15 and a light chain of the anti IL-7R antibody has the amino acid sequence shown in SEQ ID NO. 16;
- the protein formulation has a protein concentration of between 110 and 130 g/l, preferably of about 120 g/l;
- 20 - the second excipient in the protein formulation is sucrose at a concentration of between 42 and 58 g/l, preferably of about 50 g/l;
- the final excipients include polysorbate 80 which, in the protein formulation, has a concentration of between 0.017 and 0.023 g/l, preferably of about 0.02 g/l;
- 25 - the final excipients include EDTA which, in the protein formulation, has a concentration of between 0.42 and 0.58 g/l, preferably of about 0.5 g/l;
- the final excipients include arginine which, in the protein formulation, has a concentration of between 85 and 115 mM, preferably of about 100 mM;
- the protein formulation has a pH of between 6.5 and 7.5, preferably of about 30 7.0;
- the solution provided in step (a) has a protein concentration of between 2.6 and 3.4 g/l, preferably of about 3 g/l;
- the protein is concentrated to between 36 and 54 g/l, preferably to between 40 and 50 g/l, and most preferably to about 45 g/l, by the first ultra-filtration step;

- the protein is concentrated to between 170 and 210 g/l, preferably to about 190 g/l, by the second ultra-filtration step;
- the first excipient in the diafiltration buffer has a concentration higher than the concentration of the first excipient in the protein formulation, said concentration of the first excipient in the diafiltration buffer being preferably of between 19 and 25 mM, most preferably of about 22 mM;
- the diafiltration buffer includes arginine at a concentration of between 95 and 125 mM, preferably of about 110 mM;
- the diafiltration buffer has a pH of between 6.5 and 7.5, preferably about 7.0;
- adding the second excipient to the retentate obtained from the diafiltration step is achieved by adding a first additive solution to the retentate, said first additive solution comprising the second excipient at a concentration of between 230 and 320 g/l, preferably of about 275 g/l;
- the first additive solution comprises the first excipient at a concentration substantially equal to the concentration of the first excipient in the diafiltration buffer and higher than the concentration of the first excipient in the protein formulation, said concentration of the first excipient in the first additive solution being preferably of between 19 and 25 mM, most preferably of about 22 mM;
- the first additive solution further comprises a final excipient;
- the first additive solution comprises about 22 mM histidine, 110 mM arginine and about 275 g/l sucrose, at a pH of about 7.0;
- adding the first additive solution to the retentate is performed at a dilution ratio of about 5, whereby one volume of the first additive solution is added to approximately 4 fold the same volume of the retentate;
- adding the final excipients includes the step of adding a second additive solution to the solution obtained from the second ultra-filtration step, said second additive solution comprising EDTA and polysorbate 80;
- adding the second additive solution is performed at a dilution ratio of about 20, whereby one volume of the second additive solution is added to approximately 19 fold the same volume of to the solution obtained from the second ultra-filtration step.

According to a second aspect of the invention, there is provided an antibody formulation produced by the foregoing method.

- 5 In a preferred embodiment, the protein formulation comprises:
 - from 135 mg/ml to 165 mg/ml, preferably about 150 mg/ml, of the anti-PCSK9 antibody, and
 - from 16 mM to 24 mM, preferably about 20 mM, of histidine buffer.
- 10 In another preferred embodiment, the protein formulation comprises:
 - from 135 mg/ml to 165 mg/ml, preferably about 150 mg/ml, of the anti-PCSK9 antibody, and
 - from 67.2 mg/ml to 100.8 mg/ml, preferably about 84 mg/ml, of trehalose.
- 15 In another preferred embodiment, the protein formulation comprises:
 - from 135 mg/ml to 165 mg/ml, preferably about 150 mg/ml, of the anti-PCSK9 antibody, and
 - from 0.16 mg/ml to 0.24 mg/ml, preferably about 0.2 mg/ml, of polysorbate.
- 20 In another preferred embodiment, the protein formulation comprises:
 - from 135 mg/ml to 165 mg/ml, preferably about 150 mg/ml, of the anti-PCSK9 antibody,
 - from 16 mM to 24 mM, preferably about 20 mM, of histidine buffer, and
 - from 67.2 mg/ml to 100.8 mg/ml, preferably about 84 mg/ml, of trehalose.
- 25 In another preferred embodiment, the protein formulation comprises:
 - from 135 mg/ml to 165 mg/ml, preferably about 150 mg/ml, of the anti-PCSK9 antibody,
 - from 16 mM to 24 mM, preferably about 20 mM, of histidine buffer, and
 - from 0.16 mg/ml to 0.24 mg/ml, preferably about 0.2 mg/ml, of polysorbate.
- 30 In another preferred embodiment, the protein formulation comprises:
 - from 135 mg/ml to 165 mg/ml, preferably about 150 mg/ml, of the anti-PCSK9 antibody,

- from 67.2 mg/ml to 100.8 mg/ml, preferably about 84 mg/ml, of trehalose, and
- from 0.16 mg/ml to 0.24 mg/ml, preferably about 0.2 mg/ml, of polysorbate.

5 In a still preferred embodiment, the protein formulation comprises:

- from 135 mg/ml to 165 mg/ml, preferably about 150 mg/ml, of the anti-PCSK9 antibody,
- from 16 mM to 24 mM, preferably about 20 mM, of histidine buffer,
- from 67.2 mg/ml to 100.8 mg/ml, preferably about 84 mg/ml, of trehalose, and
- 10 • from 0.16 mg/ml to 0.24 mg/ml, preferably about 0.2 mg/ml, of polysorbate.

In some embodiments, the antibody formulation has a pH of between 5.2 and 5.8, preferably about 5.5.

15 In another preferred embodiment, the protein formulation comprises:

- from 110 g/l to 130 g/l, preferably about 120 g/l, of the anti-IL-7R antibody;
- from 17 mM to 23 mM, preferably about 20 mM, of histidine;
- from 42 g/l to 58 g/l, preferably about 50 g/l, of sucrose; and
- from 0.017 g/l to 0.023 g/l, preferably about 0.02 g/l, of polysorbate

20 and has a pH of between 6.5 and 7.5, preferably about 7.0.

SEQ ID NO: 1 to 12 referred to in the foregoing are described in the table below:

1 (VH)	<u>qvqlvqsgae</u> <u>vkpgasvkv</u> <u>sckasgytft</u> <u>syymhwrrqa</u> <u>pgqglewmge</u> <u>ispfggrtny</u> <u>nekfksrvtm</u> <u>trdtststvy</u> <u>melsslrned</u> <u>tavyycarer</u> <u>plyasdlwqq</u> <u>gttvvss</u>
2 (VL)	<u>digmtqspss</u> <u>lsasvgdrvt</u> <u>itcrasqqis</u> <u>salawyqqkp</u> <u>gkapkliys</u> <u>asyrytgvps</u> <u>rfsgsgsgtq</u> <u>ftftisslqp</u> <u>editatyyccqq</u> <u>ryslwrtfgq</u> <u>gkleik</u>
3 (VH – CDR1)	SYYMH
4 (VH – CDR1)	GYTFTSY
5 (VH-CDR1)	GYTFTSY ^Y MH
6 (VH-CDR2)	EISPFGGRTNYNEKFKS
7 (VH-CDR2)	ISPFGGR
8 (VH-CDR3)	ERPLYASDL
9 (VL-CDR1)	RASQGISSALA
10 (VL-CDR2)	SASYRYT

11 (VL-CDR3)	QQRYSLWRT
12 (PCSK9 human)	MGTVSSRRSW WPLPLLLLLL LLLGPAGARA QEDEDGDYEE LVLALRSEED GLAEAPEHGT TATFHRCAKD PWRLPGTYVV VLKEETHLSQ SERTARRLQA QAARRGYLTK ILHVFHGLLP GFLVKMSGDL LELALKLPHV DYIEEDSSVF AQSIPWNLER ITPPRYRADE YQPPDGGSLEVYVLLTSIQ SDHREIEGRV MVTDFENVPE EDGTRFHRQA SKCDSHGTHL AGVVSGRDAG VAKGASMRSI RVLNCQGKGT VSGTLIGLEF IRKSQQLVQPV GPLVVLPLA GGYSRVLNAA CQRLARAGVV LVTAAGNFRD DACLYSPASA PEVITVGATN AQDQPVTLGT LGTNFGRCVD LFAPGEDIIG ASSDCSTCFV SQSGTSQAAA HVAGIAAMML SAEPELTIAE LRQRLLIHSFAKDVINEAWFP EDQRVLTPNL VAALPPSTHG AGWQLFCRTV WSAHSGPTRM ATAVARCAPD EELLSCSSFS RSGKRRGERM EAQGGKLVCR AHNAFGGEGV YAIARCCLLP QANCSVHTAPPAEASMGTRV HCHQQGHVLT GCSSHWEVED LGTHKPPVLR PRGQPNCVG HREASIHASC CHAPGLECKV KEHGIPAPQE QVTVACEEGW TLTGCSALPG TSHVLGAYAV DNTCVVRSRDVSTTGSTSEG AVTAVAICCR SRHLAQASQE LQ

SEQ ID Nos. 13-16 in the forgoing are described in the table below

13 (VH)	EVQLVESGGGLVKPGGSLRLSCAASGFTFDDSVMHWRQAPGKGLEWWSLV GWDGFFTYYADSVKGRFTISRDNAKNSLYLQMNSLRAEDTAVYYCARQGDYM GNNWGQGTIVTVSS
14 (VL)	NFML TQPHSVSESPGKTVTISCTRSSGSIDSSYVQWYQQRPGSSPTTVI YEDDQRPSGVPDFRSGSIDSSNSASLTISGLKTEDEADYYCQSYDFHH LVFGGGTKLTVL
15	EVQLVESGGGLVKPGGSLRLSCAASGFTFDDSVMHWRQAPGKGLEW VSLVGWDGFFTYYADSVKGRFTISRDNAKNSLYLQMNSLRAEDTAVYYC ARQGDYMGNNWGQGTIVTVSSASTKGPSVFPLAPSSKSTSGGTAALG CLVKDVFPEPVTVSWNSGALTSGVHTFPAVLQSSGLYSLSSVTVPSSS LGTQTYICNVNHPKSNTKVDKKVAPELLGGPSVFLFPPKPKDTLMISRTP EVTCVVVDVSHEDPEVKFNWYVDGVEVHNAKTPREEQYNSTYRVVSV LTVLHQDWLNGKEYKCKVSNKALPAPIEKTIKAKGQPREPQVYTLPPSR EEMTKNQVSL TCLVKGFYPSDIAVEWESNGQPENNYKTPPVLDSDGSF FL YSKL TVDKSRWQQGNVFSCSVMHEALHNHYTQKSLSLSPGK
16	NFML TQPHSVSESPGKTVTISCTRSSGSIDSSYVQWYQQRPGSSPTTVI YEDDQRPSGVPDFRSGSIDSSNSASLTISGLKTEDEADYYCQSYDFHH LVFGGGTKLTVLQPKAAPSVTLFPPSSEELQANKATLVCISDFYPGAVT VAWKADSSPVKAGVETTPSKQSNNKYAASSYSLTPEQWKSHRSYSC QVTHEGSTVEKTVAPTECS
17 (VH-CDR1)	DSVMH
18 (VH-CDR2)	LVGWDGFFTYYADSVKG
19 (VH-CDR3)	QGDYMGNN
20 (VL-CDR1)	TRSSGSIDSSYVQ

21 (VL-CDR2)	EDDQRPS
22 (VL-CDR3)	QSYDFHHLV

Detailed description

5 The following definitions will be used in the present description and claims:

- the term "protein formulation" designates the final product including the protein of interest and excipients. When referring to proteins intended for a therapeutic use, the term "Drug Substance" may be used instead of "protein formulation" and the protein of interest may be designated by the term "active ingredient" or "product". The "excipients" are defined by all the constituents of the "protein formulation", which are not the "protein" or "active ingredient". The excipients typically include protein stabilizers, surfactants, amino-acids e.g. contributing to protein stabilization, etc...;
- in connection with the dia-filtration step, the term "retentate" refers to the solution retained on the retentate side of the membrane and containing the molecules that are too large to pass through the membrane, such as the protein of interest. The retentate is the solution that is transferred to the subsequent part of the ultra-filtration / dia-filtration system. The other solution circulated on the other side (the permeate side) of the membrane in the dia-filtration part of the system is referred to as the "dia-filtration buffer" (or "basal buffer");
- the term "concentrated pool" designates the solution directly obtained from the final ultra-filtration step;
- the term "final excipients" designates the excipients that are added to the "concentrated pool" after the final ultra-filtration step i.e. after the final concentration step;
- the term "nx spike", with n a numeral value, designates a solution of excipients that is added to a certain volume of the protein-containing solution, with a dilution ratio equal to n, which means that one volume of the solution of

excipients is added to n-1 fold the same volume of the protein-containing solution. For example a 4x spike is a solution that is added according to the ratio: 1 volume of the spike for 3 volumes of the protein-containing solution;

- 5 - unless stated otherwise, the terms "approximately", "about" or "substantially" associated with a numeral value mean within a range of $\pm 5\%$ of said value;
- "viscosity," as used herein, may be "absolute viscosity" or "kinematic viscosity." "Absolute viscosity," sometimes called dynamic or simple viscosity, is a quantity that describes a fluid's resistance to flow. "Kinematic viscosity" is

10 the quotient of absolute viscosity and fluid density. Kinematic viscosity is frequently reported when characterizing the resistive flow of a fluid using a capillary viscometer. When two fluids of equal volume are placed in identical capillary viscometers and allowed to flow by gravity, a viscous fluid takes longer than a less viscous fluid to flow through the capillary. If one fluid takes

15 200 seconds to complete its flow and another fluid takes 400 seconds, the second fluid is twice as viscous as the first on a kinematic viscosity scale. If both fluids have equal density, the second fluid is twice as viscous as the first on an absolute viscosity scale. The dimensions of kinematic viscosity are L^2/T where L represents length and T represents time. The SI units of kinematic viscosity are m^2/s . Commonly, kinematic viscosity is expressed in centistokes, cSt, which is equivalent to mm^2/s . The dimensions of absolute viscosity are $M/L/T$, where M represents mass and L and T represent length and time, respectively. The SI units of absolute viscosity are $Pa\cdot s$, which is equivalent to $kg/m\cdot s$. The absolute viscosity is commonly expressed in units of centiPoise, cP, which is equivalent to milliPascal-second, $mPa\cdot s$. In the context of the invention, an antibody is deemed to be of high viscosity if its viscosity is at

20 25 least 20 cP.

For conciseness, the acronyms "UF", "DF" and "UF/DF" (or "UFDF") may be used across the description and should be understood as follows: "UF" means "ultra-filtration, "DF" means "dia-filtration" and "UF/DF" (or "UFDF") means "ultra-filtration / dia-filtration". The method of the invention, which is defined as a method of preparation of a protein formulation, may be referred to as a UFDF method.

The invention will now be further illustrated by the following Examples, each in connection with a specific therapeutic monoclonal antibody and a specific formulation of this monoclonal antibody. The Examples are provided for illustrative 5 purpose only and should not be construed as limiting the scope of the invention.

A – Example 1

In illustrative Example 1, the protein of interest is bococizumab, a PCSK9-targeting 10 monoclonal antibody that specifically binds to PCSK9 (Proprotein Convertase Subtilisin Kexin type 9), e.g. SEQ ID NO: 12 or Uniprot Accession Number Q8NBP7. The method has been designed to achieve a targeted product concentration of 150 g/l in the Drug Substance, with the Drug Substance including the following excipients at a pH of 5.5:

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- histidine at a 20 mM concentration,
- trehalose at a 84 g/l concentration, and
- PS80 (PolySorbate 80) at a 0.2 g/l concentration.

It is deemed acceptable that the above requirements are achieved with a tolerance 20 of \pm 8 g/l in the protein concentration, of \pm 15% in the excipients concentration and of \pm 0.2 in the pH value.

In terms of yield, the method is required to achieve a product recovery of more than 90%.

25

Experiments have been conducted for defining preferred operating modes and establishing that the method of the invention is suitable for achieving the above requirements (while conventional methods are not). Some of these experiments are presented in the following part of the description.

30

A.1 Materials

The starting material used for experiments was a fully purified bococizumab solution that had been processed through a MabSelect® column to remove excipient

components prior to use. After MabSelect® purification, the eluate was adjusted to pH 5.0 by acetic acid, resulting in a product concentration of 17.09 g/l.

5 *Ultra-filtration / Dia-filtration Device*

All experiments were performed using a GE Crossflow® system (300 ml reservoir) fitted with Pellicon® 3 (30 KDa, C-screen, 88 cm²) regenerated cellulose membranes or Sartocon® (30 KDa E-channel 200 cm²) regenerated cellulose membranes.

10 TransMembrane Pressure (TMP) was maintained at approximately 14-22 psi with P_{Feed} less than 55 psi. Unless otherwise specified, all rinses were generated by recirculating rinse buffer for at least 15 minutes, then concentrating to the minimal working volume of the system.

15 **A.2 Experimental Design and Results**

Determination of Trehalose Solubility

An initial experiment was completed to evaluate the limit of the trehalose solubility in 30 mM histidine pH 5.35 solution (the histidine concentration and pH are adjusted from the final specifications to account for the exclusion of the histidine ion as the protein concentration increases). To obtain the 150 g/l final Drug Substance target, the minimal concentration of the concentrated pool would need to be 180 g/l with a 6x trehalose /EDTA/PS80 spike, or 187.5 g/l with a 5x trehalose/EDTA/PS80 spike.

25 At the trehalose concentration required to provide a 6x spike (~500 g/l), the trehalose did not dissolve (particulates were still present) at room temperature (22 °C) after extended stirring and had to be heated to 30 °C to dissolve. The solution was filtered through a 0.22 µm Pall Acrodisc® syringe filter under 15 psi pressure without re-precipitating at room temperature.

30 However, this manufacturing method may be difficult to scale up, therefore the maximum practical concentration for the trehalose spike may be capped at 5x (~420 g/l trehalose).

Accordingly, in a preferred process, the trehalose concentration of the spike solution may be about 400 g/l.

5 *Process Development*

A first experiment was designed to test the histidine concentration needed in the diafiltration solution, to check the histidine concentration in the dia-filtered solution at different protein concentrations (76.6 g/l and 114 g/l), and to generate material for 10 density measurement. The starting material was concentrated to 76.6 g/l using a 200 cm² Sartocon® E-channel membrane at a load capacity of 345 g/m², and then dia-filtered with 35 mM histidine, pH 5.26 buffer. The flux of the dia-filtration was 17 LMH (liters/m²/hour) at 300 LMH feed flowrate and 22 psi TMP. The material was then further concentrated to 213 g/l (data not shown) and samples of both the 15 diafiltered pool and final concentrated material were analyzed for histidine and trehalose concentration (see Table 1).

A second experiment was performed to determine if diafiltered material containing trehalose resulted in a lower final concentration versus material without trehalose in 20 the diafiltration buffer. The starting material was concentrated to 114 g/L and diafiltered with 35 mM histidine, pH 5.26 buffer. The diafiltration flux was 10 LMH under the operational conditions described in Table 2, Experiment 2A. The diafiltered solution was concentrated to 184.9 g/L at < 55 psi of feed pressure and 22 psi of TMP. The concentrated material was drained from the reservoir and combined 25 with the 35 mM histidine, pH 5.26 rinse solution to achieve a concentration of 153.7 g/L. The pool was spiked with 4x trehalose excipient buffer (30 mM histidine, 400 g/L trehalose, pH 5.4) to achieve a final protein concentration of 114 g/L. The spiked solution was then concentrated to 202.4 g/L under the operational conditions described in Table 2, Experiment 2B. The concentration step was stopped at 15 30 LMH feed flow rate due to pump limitations.

Table 1 shows that both the histidine and trehalose concentrations in all concentrated samples were within 10% of the final target specification, 20 mM histidine and 84 g/L trehalose.

This information provides an acceptable operating range of the diafiltration concentration from 75-114 g/L, within which the final excipient concentrations meet concentration specifications.

5

Table 1. Initial Evaluation Excipient Concentration Results

Sample Name	Concentration (g/L)	Histidine (mM)	Trehalose (g/L)
Diafiltration Exp 1	76.6	29.81	Not Tested
Concentration Exp 1	213	19.63	
Load Exp 2B	114	26.19	83.82
Concentration Exp 2B	202.4	18.67	76.37
Diafiltration Buffer	N/A	34.34	Not Tested

Table 2. Initial Evaluation Process Data

UFDF System and Load Charge	706263-18-Exp2A	706263-18-Exp2B
Membrane Type	Sartocon E-channel	
Membrane Area, m ²	0.02	0.02
UF Protein Challenge (g/m ²)	572	490
Concentration 1		
Load Volume (ml)	338.2	NA
Load Concentration (g/L)	33.8	
Concentration 1 (g/L)	114	
Concentration 1 Process Time (hr)	Not recorded	
Diafiltration		
Diafiltration Buffer	35 mM Histidine pH 5.26	NA
Diavolumes (TOV)	8	
Feed Flow Rate (LMH)	300	
Average Permeate Flux (LMH)	10	
Average TMP (psi)	22	
Diafiltration Process Time (hr)	4	
Concentration 2		
Spike Solution	NA	30 mM Histidine, 400 g/L trehalose pH 5.4
Post Spike Concentration (g/L)	NA	114
Average TMP (psi)	22	22
Final Flow Rate (LMH)	Not recorded	15
Product Volume (ml)	48.1	Not recorded
Product Concentration (g/L)	184.9	202.4
Concentration 2 Process Time (hr)*	Not recorded	~1
Retentate pH	5.50	5.51
Rinse		
Rinse Buffer	35 mM Histidine pH 5.26	20 mM Histidine, 84 g/L trehalose pH 5.5
Rinse Volume (ml)	26	Not recorded
Rinse Concentration (g/L)	58.7	54.5
Rinse pH	Not recorded	5.52

* The actual time was not recorded or could not be retrieved, it is based on a calculation of the flux and volume processed.

5

Additional experimentation was performed to evaluate changes in histidine and trehalose concentration as a function of protein concentration at end of the concentration 2 step. The starting material was concentrated to 105.9 g/l and diafiltered with 35 mM histidine, pH 5.29 buffer using a 200 cm² Sartocon® E-channel

10 membrane. The flux of the dia-filtration was 12 LMH at a feed flow rate of 300 LMH and 22 psi TMP. The dia-filtered material was then spiked with 4x trehalose

excipient solution. The spike solution was added directly into the reservoir and mixed for 15 minutes, then the material was concentrated to 172, 188 and 209 g/l final concentration (see Table 3). As shown in Table 4, the histidine concentration 5 dropped as the protein concentration increased, but all values were within 10% of the target concentration of 20 mM histidine, 84 g/L trehalose.

Table 3. Additional Development Process Data

UFDF System and Load Charge	Notebook: 706263-20
Membrane Type	Sartocon Slice E-channel
Membrane Area, m ²	0.02
UF Protein Challenge (g/m ²)	477
Concentration 1	
Load Volume (ml)	558.1
Load Concentration (g/L)	17.09
Final Concentration (g/L)	105.9
Diafiltration	
Diafiltration Buffer	35 mM Histidine pH 5.29
Diavolumes (TOV)	8
Feed Flow Rate (LMH)	300
Average TMP (psi)	22
Average Permeate Flux (LMH)	12
Diafiltration Time (hr)	3
Concentration 2	
Spike Solution	30 mM Histidine, 400 g/L trehalose pH 5.22
Post Spike Concentration (g/L)	78.6
Solution pH (At Spike)	5.38
Average TMP (psi)	22
Process Time (hr)*	~1
Product volume (ml)	32.1
Product Concentration (g/L)	209
Retentate pH	5.53
Yield Recovery (%)	76.6
Rinse	
Rinse Buffer	20 mM Histidine, 84 g/L trehalose pH 5.5
Rinse Volume (ml)	31.6
Rinse Concentration (g/L)	53.2
Rinse pH	5.48
Rinse Recovery (%)	19.2

* The actual time was not recorded or could not be retrieved, it is based on a 10 calculation of the flux and volume processed.

Table 4. Additional Development Excipient Concentration Results

Sample Name	Concentration (g/L)	Histidine (mM)	Trehalose (g/L)
Diafiltration Pool	109.4	27.81	Not Tested
Concentration Load	78.6	27.57	94.02
Concentration 1	172	20.15	84.65
Concentration 2	188	19.47	84.24
Concentration 3	209	18.11	81.86
Spike Buffer	N/A	27.83	392.88

The method was scaled up to the 500 L pilot scale (Lot 12P126J603-MV-B): see

5 Table 5 for process details. 517 g of Capto Adhere® purified material was concentrated to 107 g/l using a 0.5 m² Millipore® V-screen membrane, and then dia-
filtered with 35 mM histidine, pH 5.29 buffer, a feed flow rate of 1000 LMH and a feed pressure of 40 psi. The retentate was then spiked with 4x trehalose solution (30 mM histidine, 400 g/l trehalose, pH 5.22), which was added directly into the
10 reservoir taking into account the system hold-up volume. The spiked material was then concentrated to 202 g/l, and the concentrated product removed from the system. The skid was rinsed with 20 mM histidine, 84 g/l trehalose, pH 5.50 buffer, and the rinse added to the concentrated material. The measured concentration of the final combined solution was 160 g/l with an overall yield of 97.1%.

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Table 6 summarizes the excipient concentration and product quality results for the experiment, which shows that the final combined pool levels were within 10% of the aforementioned targeted concentrations, without any significant effect on product quality as measured by SEC when compared to past final UF values.

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Table 5. Pilot Scale Process Data

UFDF System and Load Charge	12P120J603-MV-B
Equipment	Millipore System
Membrane Type	Millipore 30K V screen RC
Membrane Area, m²	0.5
UF Protein Challenge (g/m²)	1142
Concentration 1	
Load Volume (L)	7.5
Load Concentration (g/L)	67.85
Process Time (minutes)	18
Final Concentration (g/L)	107
Diafiltration	
Diafiltration Buffer	35 mM histidine pH 5.3
Diavolumes (TOV)	8
Feed Flow Rate (LMH)	~1000
Average Flux (LMH)	20
Diafiltration Time (hours)	3.75
Concentration 2	
Spike Solution	30 mM Histidine, 400 g/L trehalose pH 5.22
Average TMP (psi)	<28
Final flow Rate (LMH)	108
Process Volume (L)	2.2
Process Time (hr)	1
Product Concentration (g/L)	202
Product pH	5.44
Yield Recovery (%)	85.7
Rinse	
Rinse Buffer	20 mM histidine, 84 g/L trehalose pH 5.5
Rinse Volume (L)	1.1
Rinse concentration (g/L)	53.7
Rinse pH	5.55
Rinse Recovery (%)	11.4

Table 6. Pilot Scale Excipient and Product Quality Results

Sample Name	Concentration (mg/ml)	Histidine (mM)	Trehalose (g/L)	Total HMMS	Monomer	Total LMMS
Diafiltered Pool	107	28.28	N/A	1.0	99.0	<0.1%
Post-Spike Pool	78.6	28.04	90.35	0.8	99.2	<0.1%
Concentration 2 Pool	202	21.57	88.21	0.9	99.1	<0.1%
Rinse Pool	53.7	18.94	80.47	0.7	99.2	<0.1%
Final Pool	160	21.28	88.25	1.2	98.8	<0.1%
DF Buffer	N/A	34.80	N/A	N/A		
Excipient Buffer		31.52	413.75			
Rinse Buffer		19.93	83.26			

5 *Evaluation of Dia-filtration process*

Protein Density and Viscosity at Different Concentrations

Figure 1 plots the viscosity of bococizumab versus product concentration in (i) 20 mM histidine, pH 5.5 and (ii) 20 mM histidine, 84 g/l trehalose, pH 5.5. The graph shows that at approximately 175 g/l the viscosity reaches the 30 cP value, which is considered the cutoff for viable UFDF processing at large scale.

Densities of bococizumab in (i) 20 mM histidine, pH 5.5 and (ii) 20 mM histidine, 84 g/l trehalose, pH 5.5 solutions were measured and are shown in Figure 2 and Figure 3. The data shows that the density is slightly less in histidine buffer as compared to histidine/trehalose buffer, which is as expected.

Based on the experiments above, it was found that the targeted concentrations in the 20 Drug Substance could be achieved at a manufacturing scale by with a DF buffer containing histidine, without trehalose.

The results from the experiments showed not only that the method of the invention resulted in acceptable yield, protein and excipient final concentrations, but also that it required lower protein concentration prior to the excipient spike, as compared to conventional methods (158 g/L versus 188 g/L). Such a lower protein concentration

is easier to achieve on a regular basis as the process is scaled up. In addition, the method of the invention is advantageous over the conventional methods due to the better cost-of-goods profile achieved by removing trehalose from the diafiltration buffer.

It has been found that the UFDF process utilizing a Millipore® C-screen membrane, as an alternative to the Millipore® V-screen membrane, consistently resulted in a concentration greater than 175 g/l, which was sufficient to allow addition of the wash pool (rinse) while still remaining above the 158 g/l needed prior to a 20x excipient spike. To ensure that the process would work with a trehalose spike, the process utilizing a C-Screen membrane was evaluated both at the laboratory and pilot scale.

The process was evaluated at the laboratory scale using Millipore® PLCTK C-Screen cassettes, utilizing the ultrafiltration (UF) run conditions outlined in Table 7.

For the TFF (Tangential flow Filtration) equipment, the lab scale process was performed employing a feed flow rate range of 30-300 LMH at an achievable pressure limit of approximately 50-55 psi as the operational limits. The upper feed flow rate of 300 LMH, where most of the process will occur, has a principle impact on process time, where reduced feed flow results in lower process flux which increases process pump time. The lower feed flow rate of 30 LMH is critical to the final concentration achievable, due to the increased viscosity increasing the pressure drop through the retentate channels, therefore lower flow rates enable pumping of more viscous solutions.

During the lab scale process run in the presence of approximately 84 g/l Trehalose a final concentration of 177 g/l was achieved in the final retentate pool with a feed flow rate of 30 LMH and feed pressure of 50 psi. The wash fractions from the lab scale runs were measured separately for yield as displayed in Table 8.

Table 7. Laboratory Scale Run Conditions

Step	Solution	Feed Pressure (psig)	Retentate Pressure (psig)	Target
Equilibration	10 mM Histidine, 50 mM NaCl, pH 6.4	20 (\pm 2)	10 (\pm 2)	\pm 0.2 pH Units
Concentration 1	VRF Product Pool	34 (\pm 6)	16 (\pm 6)	600-1000 g/m ²
Diafiltration	35 mM Histidine pH 5.3	34 (\pm 6)	16 (\pm 6)	> 7 TOV
Concentration 2	Diafiltration Pool	35 (\pm 20)	10 (\pm 10)	20-30% over DS Target
Buffer Flush	20 mM Histidine, 84 g/L Trehalose, pH 5.5	30 (\pm 20)	10 (\pm 10)	Concentration Dependent

Table 8. Results of Laboratory Scale Development Experiments

Experiment	Load Challenge (g/m ²)	Conc. 1 Flux (LMH) & Pressures (P _{feed} /P _{ret})	Diafiltration Flux (LMH) & Pressures (P _{feed} /P _{ret})	Conc. 2 Flux (LMH) & Pressures (P _{feed} /P _{ret})	% Yield Ret / Flush
No Trehalose	354*	90-20 (35/15)	25-42 (35/15)	35-4 (50/0)	85 /13
With Trehalose	354*	20-90 (35/15)	25-42 (35/15)	45-4 (50/0)	85 /13

5 *Lower load challenge is employed at lab scale due to material limitations and process cycling time and represents a worst case yield recovery option.

The laboratory scale process flux profile may be seen in Figure 4, where the vertical line indicates the starting point for reducing the feed flow rate to keep the feed 10 pressure below ~50 psi with an open retentate (zero psi), where a reduction in process flux occurs due to reducing the cross flow rate. Figure 5 shows the process feed channel pressure drop and the feed flow rate during the final concentration. In the lab scale system, the flow is manually adjusted by reducing the feed pump rate as the feed pressure approaches ~50 psi.

15

Pilot Scale Confirmation Batches

The UFDF process with the C-screen membrane was performed at the 500 L scale to confirm that the final concentration targets could be achieved. The process is

shown in Table 9 and the process data for the 3 lots performed in the pilot facility is shown in Table 10.

The results show that the process achieved high recoveries and that concentrations met the intermediate and final targets. In addition, the excipient concentrations for
5 Lot 13P120J604 were measured at 21.2 mM histidine, 85.4 g/l trehalose, and 0.051 g/l EDTA, which are within the target specifications of \pm 15%.

Table 9. UFDF Process Parameters for Pilot Scale Manufacturing

Parameter	Details	Category
Membrane	Millipore PLCTK (30 kDa cellulose) C-Screen	
Membrane Surface Area	500–1350 g/m ²	Control Limit
Operating Temperature	18-25 °C	Control Limit
Equilibration Buffer	10 mM Histidine, 50 mM NaCl, pH 6.4	
Pre-filter	≤3000 L/m ² for a 0.2 um filter	Target
Diafiltration Buffer	35 mM Histidine pH 5.3	
Dilution Buffer	30 mM Histidine, 400 g/L Trehalose, pH 5.4	
Final UF Flush Buffer	20 mM Histidine, 84 g/L Trehalose, pH 5.5	
Filter Conditioning / Equilibration	≥ 10 L/m ²	Target Range
Maximum Inlet & Retentate Pressure	≤ 80 psig	Control Limit
Concentration 1 Inlet Pressure	Setpoint 35 psig, target range 22-55 psig	Target Range
Concentration 1 Retentate Pressure	Setpoint 15 psig, target range 0-40 psig	Target Range
Target Crossflow Rate	0–10 L/min/m ²	Target Range
Permeate Flux	0–50 LMH	Expected
Diafiltration Inlet Pressure	Setpoint 40 psig, target range 22-55 psig	Target Range
Diafiltration Retentate Pressure	Setpoint 15 psig, target range 0-40 psig	Target Range
Diafiltration Concentration	70-90 g/L based on tank volume of 350-450 L	Target
	70–110 g/L	Target Range
Diafiltration Volume	Minimum 8 TOV's	Target Range
Permeate pH and Conductivity (Diafiltration End)	After 8 TOVs verify pH is 5.5 ± 0.20 units. Continue diafiltration until target is met.	Control Limit
4X Trehalose Buffer Addition	Add 30 mM Histidine, 400 g/L Trehalose pH 5.4 after DF at 1:3 ratio	Control Limit
Concentration 2 Inlet Pressure	Setpoint 22-55 psig, target range 20-60 psig	Target Range
Concentration 2 Retentate Pressure	Setpoint 0 psig, target range 0-40 psig	Target Range
Retentate Concentration	170-190 g/L (per material balance in tank)	Target
	> 158 g/L (actual retentate concentration)	Control Limit
Buffer Flush	minimal volume recirculated rinse	
UF Pool Concentration Target	158 ± 10 g/L before 20X EDTA/PS80 Spike	Control Limit
UF Pool Density	Density calculation: 0.0004 * Concentration + 1.0126	
Post-filter	≤450 L/m ² for a 0.2 um Filter	Target
Process Notes	Diafiltration Concentration based on 32-42 kg in 350 – 450 L retentate tank volume Add 4X trehalose buffer directly into tank after diafiltration. Addition at 1:3 ratio of total volume (volume in tank + system hold volume). Circulate for 10 minutes and concentrate to >158 mg/ml.	

Table 10. UFDF Process Data for 3 Pilot Scale Batches

Batch #	UNITS	13P120J604	13P120J605	13P120J606
UF Filter Type		30kD Millipore C-screen	30kD Millipore C-screen	30kD Millipore C-screen
UF Total Area	m ²	1,14	1,14	1,14
Load Volume	L	68.9L / 30.56L	81.53L / 26.57L	85.2L / 32.0L
Load Concentration	g/L	11.03 g/L / 10.9 g/L	11.04 g/L / 11.22 g/L	8.79 g/L / 11.51 g/L
UF Protein Challenge	g/m ²	959	950	982
UF Volumetric Challenge	l/m ²	87,2	95	103
Concentration 1 Start Time	AM/PM	08:21	08:21	09:08
Concentration 1 End Time	AM/PM	11:38	12:07	15:02
Concentration 1 Permeate Volume	L	86,66	97,66	107,16
Concentration 1 Feed Pressure	psig	35	35	35
Concentration 1 Retentate Pressure	psig	15	15	15
Concentration 1 Permeate Pressure	psig	0	0	0
Concentration 1 End Retentate volume	L	12,8	10,4	10,2
Concentration 1 Average Flux	L/m ² /h	23,1	22,7	15,9
Concentration 1 Average TMP	psig	25	25	25
Diafiltration Start Time	AM/PM	11:49	12:14	10:05
Diafiltration End Time	AM/PM	14:45	16:22	13:39
Diafiltration Permeate Volume	L	109	82,76	103
DiaVolumes		8,0	8,0	8,0
Diafiltration Feed Pressure	psig	40	40	40
Diafiltration Retentate Pressure	psig	15	15	15
Diafiltration Permeate Pressure	psig	0	0	0
Diafiltration Average Flux	L/m ² /h	32,6	18,8	25,3
Diafiltration Average TMP	psig	27,5	27,5	27,5
Concentration 2 Start Time	AM/PM	15:15	16:45	14:02
Concentration 2 End Time	AM/PM	15:55	17:27	14:45
Concentration 2 Feed Pressure	psig	25 – 55	25 – 55	25 – 55
Concentration 2 Retentate Pressure	psig	0 – 2	0 – 2	0 – 2
Concentration 2 Permeate Pressure	psig	0	0	0
Concentration 2 Permeate Volume	L	7,1	7,9	6,8
Concentration 2 Average Flux	L/m ² /h	9,3	9,9	8,3

Concentration 2 Average TMP (psig)		20	20	20
Product Pool Volume	L	6,433	6,337	7,175
Product Pool Concentration	g/L	158,6	157,5	156,6
Recovery	%	95	92	99
Wash Pool Volume	L	833	915	790
Wash Pool Concentration	g/L	62,88	63,20	54,28
Wash Pool Grams	g	52	58	43
Wash Pool Ratio	g/m ²	5,14%	5,79%	3,82%

A.3 Conclusions

5 The above-described experiments were able to demonstrate 92-99% step yield while achieving the Drug Substance target.

The Example demonstrates that a UFDF method according to the invention is suitable for the preparation of a highly concentrated (150 g/l) bococizumab Drug Substance with the pH and all excipient concentrations in the acceptable ranges.

10 The same may be achieved with other proteins with the same benefits, especially with proteins having a particularly high viscosity.

B – Example 2

15

In illustrative Example 2, the protein of interest is antibody C1GM, an IL-7R antagonist monoclonal antibody that specifically binds to IL-7R. The method has been designed to achieve a targeted product concentration of 120 g/l in the Drug Substance, with the Drug Substance including the following excipients at a pH of 7.0:

20

- histidine at a 20 mM concentration,
- arginine at a 100 mM concentration,
- sucrose at a 50 g/l concentration,
- PS80 (PolySorbate 80) at a 0.02 g/l concentration, and
- EDTA at a 0.5 g/l concentration.

25

It is deemed acceptable that the above requirements are achieved with a tolerance of ± 10 g/l in the protein concentration, of $\pm 15\%$ in the excipients concentration and of ± 0.5 in the pH value.

5

In terms of yield, the method is required to achieve a product recovery of more than 85%.

10 The starting material used for the experiments described below was a fully purified solution that had been processed through MabSelect® and Q membrane chromatography.

Ultrafiltration/Diafiltration Device

15 All experiments were performed using a GE Crossflow system (300 mL reservoir) or the Quattroflow™ pump system fitted with Pellicon 3® (30 KDa, C-screen, 88 cm²) regenerated cellulose membranes. Transmembrane pressure (TMP) was maintained at approximately 14-22 psi with P_{Feed} < 55 psi. Unless otherwise specified, all rinses were generated by recirculating rinse buffer for >15 minutes, 20 then concentrating to the minimal working volume of the system.

Analytical Assays

25 UV-visible spectrophotometry for protein concentration was performed using the Thermo Scientific Nanodrop 2000C™, or Solo VPE™ from C Technologies Inc. The extinction coefficient at 280 nm, as determined experimentally by ARD, is 1.51 mL*mg⁻¹*cm⁻¹.

Experiments

30

Experiment 1

The starting material was spiked with 5% of 2 M NaCl and adjusted to pH 7.0 with 2 M Tris base, concentrated to 50 g/L, diafiltered with 22 mM histidine, 110 mM

arginine pH 7.0, spiked with 5X sucrose buffer (22 mM histidine, 110 mM arginine, 275 g/L sucrose pH 7.0), and concentrated to 146.9 g/L at a feed flow rate of ~34 LMH, as detailed in Table 11. The pH of the concentrated solution was 7.00. The

5 UF system was flushed in a single pass mode (without recirculation) with the

diafiltration solution, resulting in a concentration of 33 g/L. The overall yield was approximately 88%.

Table 11 - Diafiltration and Concentration with Arginine Buffer pH 7.0

Notebook	706263-77
Pump	Quattro Flow
UF Protein Challenge (g/m ²)	348
Concentration 1	
Load Volume (L)	1.36
Load Concentration (g/L)	2.25
Feed Flow Rate (LMH)	Varied
Feed Pressure (psi)	<50
Concentration 1 Time (minutes)	320
Concentration 1 Concentration (g/L)	49.8
Diafiltration	
Diafiltration Buffer	22 mM histidine, 110 mM Arginine, pH 7.0
DiaVolumes (TOV)	8
Feed Flow Rate @ DF (LMH)	300
Feed Pressure @ DF (psi)	40
TMP @ DF (psi)	20
Permeate Flow Rate @ DF (LMH)	11.5-16.5
Diafiltration Time (minutes)	220
Concentration 2	
Spike Solution	22 mM histidine, 110 mM Arginine, 275 g/L sucrose pH 7.0
TMP @ Concentration 2 (psi)	<25
Max Feed Pressure (psi)	50
Flow rate at End (LMH)	34
Concentration 2 Time (minutes)	Not recorded
Final Concentration (g/L)	146.9

5 Tables 12-14 show the excipient, CGE and SEC assay results. The histidine, arginine, and sucrose concentrations in the final concentrated material were all

within $\pm 10\%$ of the desired value. There was no new aggregation formed during the UF process, nor any change in the level of fragmentation.

Table 12 - Process Excipient Concentrations for Arginine Buffer pH 7.0

Sample Name	Protein Concentration (g/L)	Arginine (mM)	Histidine (mM)	Sucrose (g/L)
Diafiltrate	50.1	113.7	22.6	N/A
Post-Spike	38.2	110.0	21.9	55.0
Final Concentrate	146.9	108.5	19.8	50.0
System Rinse	33	111.1	22.1	49.9
Diafiltration Buffer	N/A	116.1	23.6	N/A
5X Sucrose Buffer	N/A	87.6	15.8	277.6

5

Table 13 - nrCGE and rCGE Results for Arginine Buffer pH 7.0 Experiment

Sample	nrCGE			rCGE		
	% IgG	% Fragment	% Other	% HC + LC	% Fragment	% Other
ANTI-IL-7R Reference	97.5	2.5	0	98.7	0.5	0.8
Diafiltrate	96.2	3.8	0	98.8	0.5	0.7
Post-Spike	96.2	3.8	0	99.1	0.3	0.5
Final Concentrate	96.3	3.7	0	98.9	0.5	0.6
System Rinse	96.8	3.2	0	99.1	0.3	0.6

Table 14 - SEC Results for Arginine Buffer pH 7.0 Experiment

Sample Name	% HMMS	% LMMS	% Monomer
Diafiltrate	0.6	0.2	99.2
Post-Spike	0.6	0.2	99.2
Final Concentrate	0.6	0.2	99.2
System Rinse	0.6	0.2	99.2

5 *Experiment 2*

The experiment was repeated, as detailed in Table 15, the only difference being the way the spike volume was calculated. The total volume in the UF system prior to the spike was calculated from the overall material balance (the total load divided by the 10 diafiltrate concentration) versus adding the volume in the reservoir plus the system hold-up volume. After the 5X sucrose spike, the protein was concentrated to 183.6 g/L at ~34 LMH feed flow rate and $P_{feed} < 50$ psi.

Table 15 - Diafiltration and Concentration with Arginine Buffer at pH 7.0

Notebook	706263-78
Pump	Quattro Flow
UF Protein Challenge (g/m ²)	387
Concentration 1	
Load Volume (L)	1.5
Load Concentration (g/L)	2.27
Feed Flow Rate (LMH)	varied
Feed Pressure (psi)	<50
Concentration 1 Time (minutes)	400
Concentration 1 Concentration (g/L)	43
Diafiltration	
Diafiltration Buffer	22 mM histidine, 110 mM Arginine, pH 7.0
DiaVolumes (TOV)	10
Feed Flow Rate @ DF (LMH)	300
Feed Pressure @ DF (psi)	28
TMP @ DF (psi)	20
Permeate Flow Rate @ DF (LMH)	13.4-19.1
Diafiltration Time (minutes)	300
Concentration 2	
Spike Solution	22 mM histidine, 110 mM Arginine, 275 g/L sucrose pH 7.0
TMP @ Concentration 2 (psi)	<25
Max Feed Pressure (psi)	50
Flow Rate at End (LMH)	34
Concentration 2 Time (minutes)	Not recorded
Final Concentration (g/L)	183.6

Table 16 shows that the excipient concentrations, histidine, arginine, and sucrose concentrations in the final material were within $\pm 10\%$ of the desired target value. The difference in how the spike volume was calculated did not appear to have any significant effect on the final excipient concentrations.

Table 16 - Excipient Concentrations for Arginine pH 7.0 Buffer

Sample Name	Protein Concentration (g/L)	Arginine (mM)	Histidine (mM)	Sucrose (g/L)
Diafiltrate	43.71	112.9	21.8	ND
Post-Spike	36.2	113.1	22.1	55.6
Final Concentrate	182	106.9	19.4	48.5
5X Spike Buffer	N/A	102.1	22.2	266.7

10 *Experiment 3*

The experiment was repeated for a third time after the final formulation was nominated, and the results are detailed in Table . After diafiltration, the addition volume of the 5X spike solution was calculated as outlined in Experiment 2. The 15 protein was concentrated to 190 g/L at ~ 34 LMH feed flow rate and $P_{feed} < 50$ psi. The UF system was rinsed with 20 mM histidine, 100 mM arginine, 50 g/L sucrose, pH 7.0 in the single pass mode. The protein concentration in the combined retentate and rinse pool was 151 g/L, resulting in an overall yield of approximately 84%.

Table 17 - Diafiltration and Concentration with Arginine Buffer at pH 7.0

Notebook	706263-79
Pump	Quattro Flow
UF Protein Challenge (g/m ²)	500
Concentration 1	
Load Volume (L)	1.96
Load Concentration (g/L)	2.25
Feed Pressure (psi)	<50
Concentration 1 Time (minutes)	Not recorded
Concentration 1 Concentration (g/L)	45.4
Diafiltration	
Diafiltration Buffer	22 mM histidine, 110 mM Arginine, pH 7.0
DiaVolumes (TOV)	8
Feed Flow Rate @ DF (LMH)	450
Feed Pressure @ DF (psi)	30
TMP @ DF (psi)	25
Permeate Flow Rate @ DF (LMH)	18
Diafiltration Time (minutes)	300
Concentration 2A	
Spike Solution	22 mM histidine, 110 mM Arginine, 275 g/L sucrose pH 7.0
TMP @ Concentration 2 (psi)	<25
Max Feed Pressure (psi)	50
Flow Rate at End (LMH)	34
Concentration 2 Time (minutes)	Not recorded
Final Concentration (g/L)	190

Table 18 summarizes the excipient concentrations, showing that the histidine, arginine, and sucrose concentrations were all within $\pm 10\%$ of the desired value. Table 19 and Table 20 indicate that no additional aggregation or fragmentation was formed during the UFDF process.

Table 18 - Excipient Concentrations for Arginine pH 7.0 Buffer

Samples	Protein Concentration (g/L)	Histidine (mM)	Arginine (mM)	Sucrose (g/L)
Diafiltrate	45.4	22	110.4	N/A
Post-Spike	37.3	22	110.9	55.5
Concentration 2	190	19.5	102	49.7
System Rinse	37.6	20.5	102.2	52.2
Final ANTI-IL-7R Material	151.6	19.9	102.7	48.9
Diafiltration Buffer	N/A	22.2	110.3	N/A
5X Spike Buffer	N/A	22.2	110.4	274.4
Rinse Buffer	N/A	20	99.1	51.8

Table 19 - nrCGE and rCGE Results for Arginine pH 7.0 Buffer

Sample	nrCGE			rCGE		
	% IgG	% Fragment	% Other	% HC + LC	% Fragment	% Other
ANTI-IL-7R Reference	97.1	2.9	<0.3	98.7	0.5	0.8
ANTI-IL-7R diafiltrate	96.2	3.8	<0.3	98.9	0.4	0.7
ANTI-IL-7R spike	96.1	3.9	<0.3	99	0.4	0.7
ANTI-IL-7R concentrate	95.2	4.5	0.3	99	0.3	0.8
ANTI-IL-7R rinse	96.1	3.9	<0.3	99	0.3	0.7
ANTI-IL-7R final	95.3	4.4	0.3	99	0.3	0.7

Table 20 - SEC Results for Arginine pH 7.0 Buffer

Sample Name	% Total HMMS	% Total LMMS	% Monomer
ANTI-IL-7R Std	0.6	0.2	99.1
ANTI-IL-7R Diafiltrate	0.5	0.3	99.3
ANTI-IL-7R Spiked	0.5	0.3	99.3
ANTI-IL-7R Con	0.8	0.3	98.9
ANTI-IL-7R Rinse	0.5	0.3	99.3
ANTI-IL-7R Final	0.7	0.3	99

The process as performed in Experiment 3 above results in acceptable concentration values for all of the excipients and the protein of interest, and does not appear to have an effect on either formation of aggregate or fragmentation. This process will be scaled up to the pilot scale to ensure that it performs as expected.

Pilot Scale UFDF Process

10

The UF process developed above (Experiment 3) was tested in the Pilot Plant using a Millipore® C-screen regenerated cellulose membrane, and using material purified from a 500 L scale bioreactor.

15

During the unit operation, detailed in Table 21, 86 L of starting material at a starting product concentration of 2.92 g/L was spiked with 5% of 2 M NaCl, then concentrated to 44.6 g/L. The material was then diafiltrated with 22 mM histidine, 110 mM arginine, pH 7.0 at a feed flow rate of approximately 150 LMH and $P_{feed} < 40$ psi. After 8 TOV diafiltration, the retentate was spiked with 22 mM histidine, 110 mM arginine, 275 g/L sucrose pH 7.0, and recirculated for 10 minutes, then concentrated to 191.4 g/L. The concentration process was stopped at 30 LMH permeate flow rate and $P_{feed} < 50$ psi. The skid was then rinsed with 20 mM histidine, 100 mM histidine, 50 g/L sucrose, pH 7.0 in single pass mode. The overall yield was approximately 87%. The entire UF process took approximately 5 hours to complete.

25

A UF pool at a concentration of 135.4 g/L was created by mixing the retentate pool, the rinse pool, and additional rinse buffer. The pool was filtered through a Millipore® 05/0.2 um Opticap Express SHC at 59 L/m² throughput. A 20X EDTA and PS80 5 excipient buffer was spiked into the UF pool to produce Drug Substance at a final concentration of 129.4 g/L.

Table 21 - Pilot Scale UF Process Data

UFDF System and Load Charge	
Membrane Type	Millipore 30K C screen RC
Membrane Area, m ²	2.28
UF Protein Challenge (g/m ²)	123
Concentration 1	
Load Volume (L)	96
Load Concentration (g/L)	2.923
Concentration 1 Time (minutes)	~120
Concentration 1 Concentration (g/L)	44.6
Diafiltration	
Diafiltration Buffer	22 mM Histidine, 110 mM Arginine pH 7.0
DiaVolumes (TOV)	8
Retentate Flow Rate @ DF (LMH)	131.6
Feed Pressure @ DF (psi)	29
TMP @ DF (psi)	24
Permeate Flow Rate @ DF (LMH)	18.4
Diafiltration Time (minutes)	80
Concentration 2	
Spike Solution	22 mM Histidine, 110 mM Arginine, 275 g/L Sucrose pH 7.0
TMP @ Concentration 2 (psi)	25
Retentate Flow Rate @ End (LMH)	30
Concentration 2 Final Volume (L)	0.738
Concentration 2 Time (minutes)	~30

Concentration 2	Concentration (g/L)	191.4
Concentration 2 Recovery (%)		50.35
Rinse		
Rinse Buffer		20 mM Histidine, 100 mM Arginine, 50 g/L Sucrose pH 7.0
Rinse Final Volume (mL)		1057
Rinse Concentration (g/L)		97.3
Rinse Recovery (%)		36.7

The excipient concentrations are summarized in Table 22, which shows that the histidine, arginine, and sucrose concentrations in the final pool were within $\pm 15\%$ of the target values. The targeted range during the process development at lab scale 5 was set at $\pm 10\%$ of the target values for all excipient concentrations, but the acceptance range at large scale was set at $\pm 15\%$ to allow for latitude during scale-up.

Table 23 and Table 24 summarize the product quality results from the run, which show that no increase in aggregation or fragmentation was detected.

10

Table 22 - Pilot Scale Run Excipient Concentration Results

Sample Name	Concentration (g/L)	Arginine (mM)	Histidine (mM)	Sucrose (g/L)
Diafiltration Pool	44.6	111.4	22.2	N/A
5X Spiked Pool	32.9	111.2	22.1	46.3
Concentration 2 Retentate	191.4	105.8	20.1	43.9
Final Rinse Pool	97.3	101.8	19.9	46.4
Drug Substance	135.4	103.9	20.2	43.8
UF Buffer	N/A	110.6	22.2	N/A
5X Spike	N/A	112.7	22.5	269.5
Rinse Buffer	N/A	101.0	20.4	46.9

Table 23 - Pilot Scale Run SEC Results

Sample Name	% Total HMMS	% Total LMMS	% Monomer
ANTI-IL-7R Reference	0.6	0.4	99.0
UF Load	0.5	0.4	99.2
Concentration 2 UF Retentate	0.7	0.4	98.9
Final Rinse Pool	0.6	0.3	99.1
Drug Substance	0.6	0.3	99.0

Table 24 - Pilot Scale Run nrCGE Results

Sample	% IgG	% Fragment	% Other
ANTI-IL-7R Reference	96.5	3.1	0.3
UF Load	96.7	3.3	0.0
Diafiltration Pool	97.3	2.7	0.0
Concentration 2 UF Retentate	96.2	3.4	0.3
Final Rinse Pool	97.1	2.9	0.0
Drug Substance	96.5	3.5	0.0

5

Conclusions

In conclusion, the above-described experiments demonstrate the successful process development of a UFDF process for >120 mg/ml drug substance for the ANTI-IL-7R antibody of interest. The UFDF process includes an initial concentration, a diafiltration, a sucrose spike prior to a final concentration, then spiking with the remaining excipients. The pH and all excipient concentrations in the developed process are in the acceptable ranges.

15

CLAIMS:

1. A method of preparing a protein formulation including excipients and at least one therapeutic protein, the method comprising the sequential steps of:

(a) providing a solution comprising said therapeutic protein;

5 (b) concentrating the therapeutic protein in the solution by a first ultra-filtration step;

(c) diafiltering the solution thus obtained with a diafiltration buffer including at least one first excipient, whereby a retentate is obtained comprising the therapeutic protein and the first excipient;

10 (d) adding a second excipient to the retentate obtained from the diafiltration step, wherein the second excipient is a sugar;

(e) further concentrating the therapeutic protein in the retentate by a second ultra-filtration step in an ultra-filtration equipment; and

15 (f) adding at least one final excipient, whereby the protein formulation with a desired protein concentration and including said first, second and final excipients is obtained,

wherein said therapeutic protein is an antibody.

2. The method of claim 1, further including, after step (e) and before step (f), rinsing the ultra-filtration equipment with a rinse buffer, whereby the recovery 20 of the protein is enhanced.

3. The method of claim 2, wherein the rinse buffer comprises the first and the second excipients at concentrations substantially equal to, respectively, the concentrations of the first and of the second excipients in the protein formulation.

4. The method of any one of claims 1 to 3, wherein the first excipient is an amino-acid.
5. The method of claim 4, wherein the first excipient is histidine.
6. The method of any one of claims 1 to 5, wherein the first excipient in the 5 protein formulation has a concentration of between 16 and 24 mM.
7. The method of claim 6, wherein the first excipient in the protein formulation has a concentration of between 17 and 23 mM.
8. The method of claim 7, wherein the first excipient in the protein formulation has a concentration of about 20 mM.
- 10 9. The method of any one of claims 1 to 8, wherein the second excipient is a disaccharide.
10. The method of any one of claims 1 to 9, wherein the final excipients include a surfactant.
11. The method of claim 10, wherein the surfactant is polysorbate 80.
- 15 12. The method of any one of claims 1 to 11, wherein the final excipients include a chelating agent.
13. The method of claim 12, wherein the chelating agent is EDTA.
14. The method of any one of claims 1 to 13, wherein the protein formulation has a protein concentration of between 110 and 165 g/l.
- 20 15. The method of any one of claims 1 to 14, wherein the antibody is an anti-IL-7R antibody.
16. The method of claim 15, wherein the antibody has a VH region comprising the amino acid sequence shown in SEQ ID NO: 13, and VL region comprising the amino acid sequence shown in SEQ ID NO: 14.

17. The method of claims 15 or 16, wherein the protein formulation has a protein concentration of between 110 and 130 g/l.

18. The method of claim 17, wherein the protein formulation has a protein concentration of about 120 g/l.

5 19. The method of any one of claims 15 to 18, wherein the second excipient in the protein formulation is sucrose at a concentration of between 42 and 58 g/l.

20. The method of claim 19, wherein the concentration of sucrose is about 50 g/l.

10 21. The method of any one of claims 15 to 20, wherein the final excipients include polysorbate 80 which, in the protein formulation, has a concentration of between 0.017 and 0.023 g/l.

22. The method of claim 21, wherein the concentration of polysorbate 80 is about 0.02 g/l.

15 23. The method of any one of claims 15 to 22, wherein the final excipients include EDTA which, in the protein formulation, has a concentration of between 0.42 and 0.58 g/l.

24. The method of claim 23, wherein the concentration of EDTA is about 0.5 g/l.

20 25. The method of any one of claims 15 to 24, wherein the final excipients include arginine which, in the protein formulation, has a concentration of between 85 and 115 mM.

26. The method of claim 25, wherein the concentration of arginine is about 100 mM.

25 27. The method of any one of claims 15 to 26, wherein the protein formulation has a pH of between 6.5 and 7.5.

28. The method of claim 27, wherein the protein formulation has a pH of about 7.0.

29. The method of any one of claims 15 to 28, wherein the solution provided in step (a) has a protein concentration of between 2.6 and 3.4 g/l.

5 30. The method of claim 29, wherein the solution provided in step (a) has a protein concentration of about 3 g/l.

31. The method of any one of claims 15 to 30, wherein the protein is concentrated to between 36 and 54 g/l by the first ultra-filtration step.

10 32. The method of claim 31, wherein the protein is concentrated to between 40 and 50 g/l by the first ultra-filtration step.

33. The method of claim 32, wherein the protein is concentrated to about 45 g/l by the first ultra-filtration step.

34. The method of any one of claims 15 to 33, wherein the protein is concentrated to between 170 and 210 g/l by the second ultra-filtration step.

15 35. The method of claim 34, wherein the protein is concentrated to about 190 g/l by the second ultra-filtration step.

36. The method of any one of claims 15 to 35, wherein the first excipient in the diafiltration buffer has a concentration higher than the concentration of the first excipient in the protein formulation.

20 37. The method of claim 36, wherein said concentration of the first excipient in the diafiltration buffer is of between 19 and 25 mM.

38. The method of claim 37, wherein said concentration of the first excipient in the diafiltration buffer is of about 22 mM.

39. The method of any one of claims 15 to 38, wherein the diafiltration buffer includes arginine at a concentration of between 95 and 125 mM.

40. The method of claim 39, wherein the diafiltration buffer includes arginine at a concentration of about 110 mM.

5 41. The method of any one of claims 15 to 40, wherein the diafiltration buffer has a pH of between 6.5 and 7.5.

42. The method of claim 41, wherein the diafiltration buffer has a pH of about 7.0.

10 43. The method of any one of claims 15 to 42, wherein adding the second excipient to the retentate obtained from the diafiltration step is achieved by adding a first additive solution to the retentate.

44. The method of claim 43, wherein said first additive solution comprises the second excipient at a concentration of between 230 and 320 g/l.

15 45. The method of claim 44, wherein said first additive solution comprises the second excipient at a concentration of about 275 g/l.

46. The method of any one of claims 43 to 45, wherein the first additive solution comprises the first excipient at a concentration substantially equal to the concentration of the first excipient in the diafiltration buffer and higher than the concentration of the first excipient in the protein formulation.

20 47. The method of claim 46, wherein said concentration of the first excipient in the first additive solution is of between 19 and 25 mM.

48. The method of claim 47, wherein said concentration of the first excipient in the first additive solution is of about 22 mM.

25 49. The method of any one of claims 43 to 48, wherein the first additive solution further comprises a final excipient.

50. The method of claim 49, wherein the first additive solution comprises about 22 mM histidine, 110 mM arginine and about 275 g/l sucrose, at a pH of about 7.0.

51. The method of any one of claims 43 to 50, wherein adding the first additive solution to the retentate is performed at a dilution ratio of about 5, whereby one volume of the first additive solution is added to approximately 4 fold the same volume of the retentate.

52. The method of any one of claims 15 to 51, wherein adding the final excipients includes the step of adding a second additive solution to the solution obtained from the second ultra-filtration step, said second additive solution comprising EDTA and polysorbate 80.

53. The method of claim 52, wherein adding the second additive solution is performed at a dilution ratio of about 20, whereby one volume of the second additive solution is added to approximately 19 fold the same volume of the solution obtained from the second ultra-filtration step.

54. A formulation of an antibody having a viscosity of at least 20 cP produced by the method of any one of claims 1 to 53.

55. The formulation of claim 54, wherein the antibody is an anti-IL-7R antibody and wherein the protein formulation comprises:

20 • from 110 g/l to 130 g/l of the anti-IL-7R antibody;

• from 17 mM to 23 mM of histidine;

• from 42 g/l to 58 g/l of sucrose; and

• from 0.017 g/l to 0.023 g/l of polysorbate

and has a pH of between 6.5 and 7.5.

56. The formulation of claim 55, wherein the antibody is an anti-IL-7R antibody and wherein the protein formulation comprises:

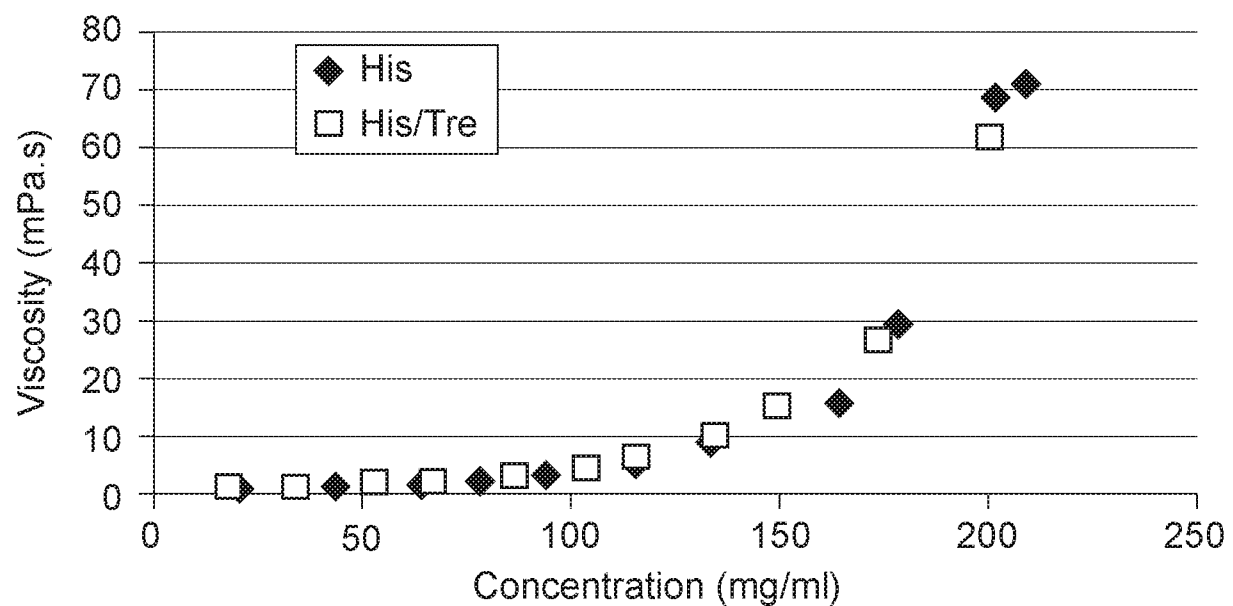
- about 120 g/l of the anti-IL-7R antibody;
- about 20 mM of histidine;
- 5 • about 50 g/l of sucrose; and
- about 0.02 g/l of polysorbate

and has a pH of about 7.0.

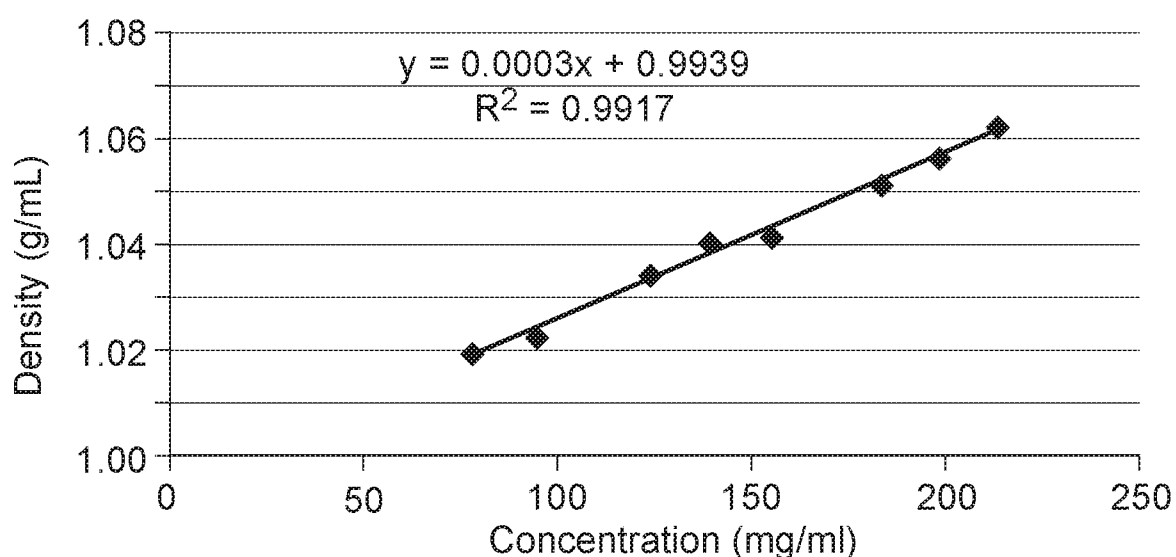
1/3

FIG. 1

Viscosity at Different Protein Concentrations in Histidine and Histidine/Trehalose

**FIG. 2**

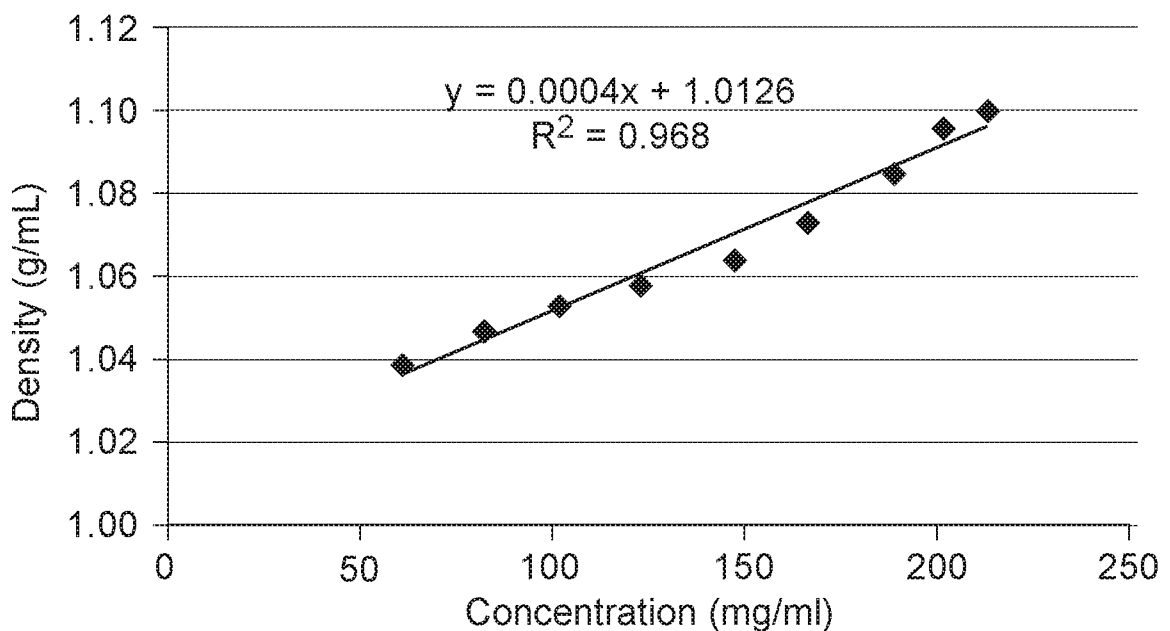
Density at Different Protein Concentrations in Histidine



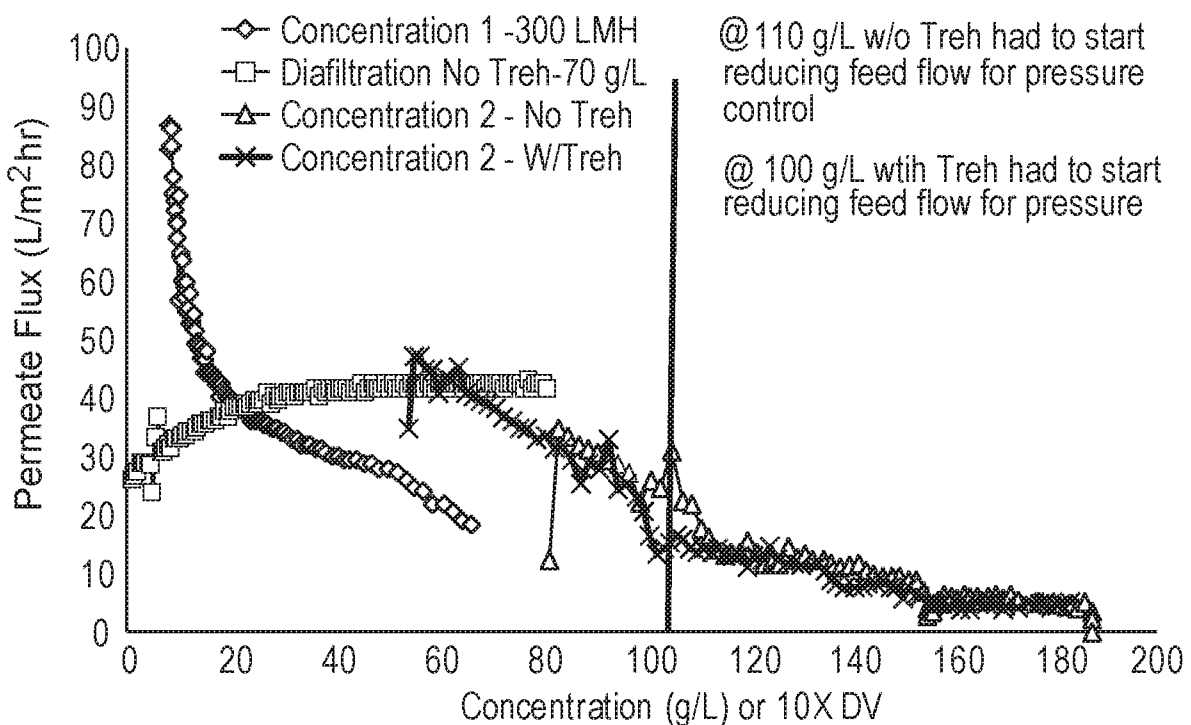
2/3

FIG. 3

Density at Different Protein Concentrations in Histidine/Trehalose

**FIG. 4**

Laboratory Scale C-Screen Process Flux



3/3

FIG. 5

Laboratory Scale C-Screen Pressure and Feed Flow Rate

